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         Jul 22
                 INPADOC: Basic index (/BI) enhanced; Simultaneous Left and
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                 Right Truncation available
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         AUG 05
                 New pricing for EUROPATFULL and PCTFULL effective
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                 August 1, 2003
         AUG 13
                 Field Availability (/FA) field enhanced in BEILSTEIN
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     9
NEWS 10
         AUG 15
                 PATDPAFULL: one FREE connect hour, per account, in
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        AUG 15
                 PCTGEN: one FREE connect hour, per account, in
NEWS 11
                 September 2003
NEWS 12
        AUG 15
                 RDISCLOSURE: one FREE connect hour, per account, in
                 September 2003
                 TEMA: one FREE connect hour, per account, in
NEWS 13
        AUG 15
                 September 2003
        AUG 18
                 Data available for download as a PDF in RDISCLOSURE
NEWS 14
NEWS 15
                 Simultaneous left and right truncation added to PASCAL
         AUG 18
                 FROSTI and KOSMET enhanced with Simultaneous Left and Righ
NEWS 16
        AUG 18
                 Truncation
        AUG 18
                 Simultaneous left and right truncation added to ANABSTR
NEWS 17
NEWS 18
        SEP 22
                 DIPPR file reloaded
NEWS 19
         SEP 25
                 INPADOC: Legal Status data to be reloaded
NEWS 20 SEP 29
                 DISSABS now available on STN
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NEWS EXPRESS
              MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP),
              AND CURRENT DISCOVER FILE IS DATED 01 APRIL 2003
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              CAS World Wide Web Site (general information)
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FILE 'HOME' ENTERED AT 15:02:18 ON 29 SEP 2003

=> file medline, uspatful, dgene, embase, wpids, biosis

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FULL ESTIMATED COST

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FILE 'MEDLINE' ENTERED AT 15:03:32 ON 29 SEP 2003

FILE 'USPATFULL' ENTERED AT 15:03:32 ON 29 SEP 2003
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=> s fermentation () xylose () ethanol L1 4 FERMENTATION (W) XYLOSE (W) ETHANOL

=> d l1 ti abs ibib tot

- L1 ANSWER 1 OF 4 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN
- TI Yeast which ferments xylose to ethanol comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated at each of its multiple reiterated ribosomal DNA sites.
- AN 1997-558974 [51] WPIDS
- AB WO 9742307 A UPAB: 19991020

Novel yeast which ferments xylose to ethanol, comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol, where the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at least 20 generations.

USE - The methods can produce yeast, which even upon culture in non-selective medium for multiple generations, e.g. up to 20, retain their full capability to ferment xylose to ethanol. Dwg.0/12

ACCESSION NUMBER:

1997-558974 [51] WPIDS

DOC. NO. CPI:

C1997-178545

TITLE:

Yeast which ferments xylose to ethanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase

genes integrated at each of its multiple reiterated

ribosomal DNA sites.

DERWENT CLASS: INVENTOR(S):

D16 D17 E17 H06 CHEN, Z; HO, N W Y

PATENT ASSIGNEE(S):

(PURD) PURDUE RES FOUND

COUNTRY COUNT:

76

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

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WO 9742307 A1 19971113 (199751) * EN 66
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RW: AT BE CH DE DK EA ES FI FR GB GH GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE HU IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG US UZ VN YU

AU 9728301 A 19971126 (199813)

EP 898616 A1 19990303 (199913) EN

R: AT BE DE DK ES FI FR GB GR IE IT NL PT SE

CN 1225125 A 19990804 (199949)

JP 2000509988 W 20000808 (200043) 50

MX 9809223 A1 19990701 (200061)

AU 731102 B 20010322 (200122)

BR 9710963 A 20010731 (200146)

APPLICATION DETAILS:

PAT	TENT NO K	IND	API	PLICATION	DATE
WO	9742307	A1	WO	1997-US7663	19970506
AU	9728301	A	ΑU	1997-28301	19970506
ΕP	898616	A1	EΡ	1997-922698	19970506
			WO	1997-US7663	19970506
CN	1225125	A	CN	1997-196195	19970506
JΡ	2000509988	W	JР	1997-540153	19970506
			WO	1997-US7663	19970506
MX	9809223	A1	MX	1998-9223	19981105
ΑU	731102	В	ΑU	1997-28301	19970506
BR	9710963	A	BR	1997-10963	19970506
			WO	1997-US7663	19970506

FILING DETAILS:

PATENT NO KIND PATENT NO									
AU 9728301	A Based on	WO 9742307							
EP 898616	Al Based on	WO 9742307							
JP 2000509988	W Based on	WO 9742307							
AU 731102	B Previous Publ.	AU 9728301							
	Based on	WO 9742307							
BR 9710963	A Based on	WO 9742307							

PRIORITY APPLN. INFO: US 1996-16865P 19960506

- L1 ANSWER 2 OF 4 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN
- Prodn. of a prod. using a bi layer pellet contg. an immobilised enzyme system used in the simultaneous isomerisation and fermentation of xylose to ethanol.
- AN 1995-122845 [16] WPIDS
- CR 1993-344962 [43]
- AB US 5397700 A UPAB: 19950502

A prod (I) is formed using bilayer pellets (outer layer of a porous polymer material (II) contg immobilised urease (III); inner core of (II) contg an immobilised enzyme (IV) other than (III) that acts on a substrate to afford (I) as follows: (a) the pellets are dispersed in a bulk soln contg urea and the substrate, and with an acidic pH; (b) (III) reacts with urea as it diffuses into the outer layer to furnish NH3 that consumes H+ diffusing into the inner core from the bulk soln to provide (IV) in the inner core with a pH higher than the acidic pH suitable for reacting with the substrate as it diffuses into the inner core; and (c) (IV) reacts with the substrate as it diffuses in to furnish (I).

ADVANTAGE - The method is partic suitable for the isomerisation of xylose to xylulose in the pellets, and the simultaneous fermentation of produced diffused xylulose (and glucose) to EtOH in the bulk soln EtOH is

a known lig fuel in gasoline additives, etc.

Dwq.0/2

ACCESSION NUMBER:

1995-122845 [16] WPIDS

CROSS REFERENCE:

1993-344962 [43]

DOC. NO. CPI:

C1995-056046

TITLE:

Prodn. of a prod. using a bi layer pellet contq. an immobilised enzyme system - used in the simultaneous isomerisation and fermentation of xylose to ethanol.

D16 E17 H06 DERWENT CLASS:

INVENTOR(S):

BYERS, J P; FOURNIER, R L; VARANASI, S

PATENT ASSIGNEE(S):

(UYTO-N) UNIV TOLEDO

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG _____

US 5397700 A 19950314 (199516)*

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
US 5397700	A CIP of	US 1991-785938 US 1993-125546	

FILING DETAILS:

PATENT NO KIND PATENT NO ______ US 5397700 A CIP of US 5254468

PRIORITY APPLN. INFO: US 1991-785938 19911031; US 1993-125546 19930923

- ANSWER 3 OF 4 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN 1.1
- Bi layer pellet contg. immobilised xylose isomerase and urease used for TIsimultaneous isomerisation and fermentation of xylose to ethanol.
- 1993-344962 [43] AN WPIDS
- 1995-122845 [16] CR
- US 5254468 A UPAB: 19950508 AΒ

Bilayered immobilised enzyme pellet comprises: (a) a core consisting of xylose isomerase (I) immobilised onto a porous polymer material (II); and (b) an outer layer consisting of urease (III) immobilised onto a porous polymer material (IV).

Pref. (IV) is polyacrylamide. The pellet is prepd. by: immobilising (I) onto (II); mixing the reulsting particles with H2O, (III), a monomer, a crosslinking agent, and a polymerisation initiator; keeping the suspension at 0-4 deg. C.; adding PhMe, CHCI3 and a surfactant; and agitating the produced aq. hydrophobic phase at 0-4 deg. C. under N2 to effect polymerisation of the monomer to form a thin polymer coating (contq. immobilised (III)) on the (I)-contq. particles.

USE/ADVANTAGE - The process allows the simultaneous isomerisation of xylose to xylulose, and the immediate fermentation of the latter cpd. to EtOH to be effected at the optimum (but different) pH values. In addn. feeds contg. xylose and glucose (i.e. as obtd. from lignocellulose) may also be used.

Dwg.0/2 Dwg.0/2

ACCESSION NUMBER:

1993-344962 [43] WPIDS

CROSS REFERENCE:

1995-122845 [16]

DOC. NO. CPI:

C1993-152813

TITLE:

Bi layer pellet contg. immobilised xylose isomerase and

urease - used for simultaneous isomerisation and

fermentation of xylose to ethanol.

DERWENT CLASS: D16 D17 E17

INVENTOR(S): BYERS, J P; FOURNIER, R L; VARANASI, S

PATENT ASSIGNEE(S): (UYTO-N) UNIV TOLEDO

COUNTRY COUNT:

PATENT INFORMATION:

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION DATE
		
US 5254468	Α	US 1991-785938 19911031

PRIORITY APPLN. INFO: US 1991-785938 19911031

L1 ANSWER 4 OF 4 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

TI Fermenting D-xylose to ethanol - using specific yeast mutants with high conversion efficiency.

AN 1982-04576J [48] WPIDS

AB WO 8204068 A UPAB: 19930915

Direct fermentation of D-xylose (I) to ethanol comprises inoculating a medium contg. nutrients and (I) with a yeast able to convert (I) to ethanol with bioconversion yield at least 50%. The mixt. is fermented until (I) conversion to ethanol of at least 50 (pref. 80)% is achieved. Pref. the yeast mutants Candida sp. XF217 or Saccharomyces cerevisiae SCXF 138 (both claimed as new microorganisms) are used. The medium contains 1-40 (5-30) wt.-vol.% (I) initially and is fermented aerobically or anaerobically at 22-40 (30) deg.C and pH 4-8 (about 6). The medium may also contain D-glucose (also converted) e.g. a cellulose or hemicellulose hydrolysate.

Hemicellulose waste materials e.g. sugar cane bagasse, are available in large quantities and then mutants efficiently convert the sugar formed when they are hydrolysed.

ABEQ US 4511656 A UPAB: 19930915

Prodn. of ethanol comprises fermentation of D-xylose with a parent yeast strain of Candida sp. or Saccharomyces cerevisiiae species, in the presence of suitable nutrients at pH about 4-8 pref. 6, and at 22-40 pref 30 deg under aerobic conditions; such that at least 50% pref. 80% of the xylose is converted to EtOH.

ADVANTAGE - Process utilises cellulose hydrolysate and/or hemicellulose hydrolysate as a nutrient medium, with conversion of both D-glucose and D-xylose.

ABEQ EP 66396 B UPAB: 19930915

A process for the direct fermentation of D-xylose to ethanol which comprises incoluating a medium comprising growth nutrients and D-xylose with a yeast mutant having an ability to ferment D-xylose to ethanol with a bioconversion yield of at least 50%, permitting the inoculated medium to ferment for a period of time sufficient to achieve a conversion of D-xylose to ethanol of at least 50% and recovering the ethanol so produced as product.

ACCESSION NUMBER: 1982-04576J [48] WPIDS

TITLE: Fermenting D-xylose to ethanol - using specific yeast

mutants with high conversion efficiency.

DERWENT CLASS: D16 D17 E17 INVENTOR(S): GONG, C S

PATENT ASSIGNEE(S): (PURD) PURDUE RES FOUND; (PURO) PUROLATOR INC

COUNTRY COUNT: 2

PATENT INFORMATION:

_____ WO 8204068 A 19821125 (198248) * EN 24 W: AU BR DK FI JP NO EP 66396 A 19821208 (198250) EN R: AT BE CH DE FR GB IT LI LU NL SE US 4368268 A 19830111 (198305) ZA 8203350 A 19830427 (198329) US 4511656 A 19850416 (198518) EP 66396 B 19850821 (198534) EN R: AT BE CH DE FR GB IT LI LU NL SE DE 3265585 G 19850926 (198540) A 19860708 (198632) CA 1207257 APPLICATION DETAILS: APPLICATION DATE PATENT NO KIND ______ EP 1982-302474 19820514 EP 66396 US 4368268 A US 1982-376731 19820511 PRIORITY APPLN. INFO: US 1981-263925 19810515; US 1981-363925 19810515; US 1982-376731 19820511 => d his (FILE 'HOME' ENTERED AT 15:02:18 ON 29 SEP 2003) FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS' ENTERED AT 15:03:32 ON 29 SEP 2003 4 S FERMENTATION () XYLOSE () ETHANOL => s yeast () xylose () ethanol 0 YEAST (W) XYLOSE (W) ETHANOL => s fermentation of xylose to ethanol 59 FERMENTATION OF XYLOSE TO ETHANOL => s genes integrated () multiple reiterated ribosomal DNA O GENES INTEGRATED (W) MULTIPLE REITERATED RIBOSOMAL DNA => s ribosomal DNA 18161 RIBOSOMAL DNA => s 15 and reiterated 58 L5 AND REITERATED => s 16 and integrated genes 0 L6 AND INTEGRATED GENES => s 17 and genes integrated 0 L7 AND GENES INTEGRATED => s 16 and genes 41 L6 AND GENES => s 19 and integrat? 12 L9 AND INTEGRAT? => d l10 ti abs ibib tot L10 ANSWER 1 OF 12 USPATFULL on STN Method for normalizing the relative intensities of detection signals in

L1

hybridization arrays

AB

The present invention relates to rRNA-derived cDNA used as an internal standard or control to achieve normalization of hybridization signal detection in microarray biochip technology. Analysis of data obtained from a laser scanner during DNA microarray experiments first requires image processing. However, the data generated for the arrayed genes must be normalized before differentially expressed genes can be identified. Normalization is necessary to compensate for differences in labelling and detection efficiencies for the labels and for differences in the quantity of starting RNA from the samples examined in the assay. Because of its relatively invariant expression across tissues and treatments, 18S and 28S ribosomal RNAs are ideal internal controls for quantitative RNA analysis. A way to circumvent the technical difficulties of using ribosomal RNA as a control, because of its overabundance relative to that of other RNAs, is described and claimed in the present application. Improved methods, arrays, and kits comprising arrays and free unlabelled ribosomal probes, are objects of this invention. The unlabelled ribosomal probes are used to compete out the excess or ribosomal nucleics present in a sample wherein all cDNA species of the sample are labelled before being placed in contact with the arrays.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:213648 USPATFULL

TITLE:

Method for normalizing the relative intensities of

detection signals in hybridization arrays

INVENTOR (S):

Larose, Anne-Marie, Montreal, CANADA LeBlanc, Benoit, Montreal, CANADA Camato, Rino, St-Leonard, CANADA

	NUMBER	KIND	DATE	
US	2003148286	A1	20030807	
US	2002-30846	A1	20020719	(10)
WO	2001-CA1860		20011221	

NUMBER DATE

PRIORITY INFORMATION:

PATENT INFORMATION: APPLICATION INFO.:

CA 2000-2327527 20001227

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE: MERCHANT & GOULD PC, P.O. BOX 2903, MINNEAPOLIS, MN,

55402-0903

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

12

NUMBER OF DRAWINGS:

6 Drawing Page(s)

LINE COUNT:

2959

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L10 ANSWER 2 OF 12 USPATFULL on STN

Identification of genes ΤT

AB A method for identifying a microorganism having a reduced adaptation to a particular environment comprising the steps of:

- (1) providing a plurality of microorganisms each of which is independently mutated by the insertional inactivation of a gene with a nucleic acid comprising a unique marker sequence so that each mutant contains a different marker sequence, or clones of the said microorganism;
- (2) providing individually a stored sample of each mutant produced by step (1) and providing individually stored nucleic acid comprising the unique marker sequence from each individual mutant;

- (3) introducing a plurality of mutants produced by step (1) into the said particular environment and allowing those microorganisms which are able to do so to grow in the said environment;
- (4) retrieving microorganisms from the said environment or a selected part thereof and isolating the nucleic acid from the retrieved microorganisms;
- (5) comparing any marker sequences in the nucleic acid isolated in step (4) to the unique marker sequence of each individual mutant stored as in step (2); and
- (6) selecting an individual mutant which does not contain any of the marker sequences as isolated in step (4).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2002:19054 USPATFULL

TITLE:

Identification of genes

INVENTOR(S):

Holden, David William, London, UNITED KINGDOM Shea, Jacqueline Elizabeth, High Wycombe, UNITED

PATENT ASSIGNEE(S):

Hensel, Michael, Munchen, GERMANY, FEDERAL REPUBLIC OF Imperial College Innovations Limited, London, UNITED

KINGDOM (non-U.S. corporation)

Microscience Limited, Berkshire, UNITED KINGDOM

(non-U.S. corporation)

			N	U	M	В	E	R							K	Ι	N	D		DATE							
_	 _	_	_	_	_	_	_	_	_	_	-	_	-	-	-	_	_	_	_	-	_	_	_	-	-	-	

PATENT INFORMATION:

US 6342215 B1 20020129

APPLICATION INFO.: RELATED APPLN. INFO.:

US 1998-201945 19981201 (9) Continuation of Ser. No. US 637759, now patented, Pat.

No. US 5876931

		NUMBER	DATE
PRIORITY	INFORMATION:	GB 1994-24921 GB 1995-1881	19941209 19950131
DOGUMENU	my DE .	GB 1995-9239	19950505

DOCUMENT TYPE: FILE SEGMENT:

Utility GRANTED

PRIMARY EXAMINER: LEGAL REPRESENTATIVE:

Schwartzman, Robert A. Holland & Knight LLP

NUMBER OF CLAIMS:

21

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

119 Drawing Figure(s); 112 Drawing Page(s)

LINE COUNT:

7399

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L10 ANSWER 3 OF 12 USPATFULL on STN
- TIIdentification of genes
- AB A method for identifying a microorganism having a reduced adaptation to a particular environment comprising the steps of:
 - (1) providing a plurality of microorganisms each of which is independently mutated by the insertional inactivation of a gene with a nucleic acid comprising a unique marker sequence so that each mutant contains a different marker sequence, or clones of the said microorganism;
 - (2) providing individually a stored sample of each mutant produced by step (1) and providing individually stored nucleic acid comprising the unique marker sequence from each individual mutant;

- (3) introducing a plurality of mutants produced by step (1) into the said particular environment and allowing those microorganisms which are able to do so to grow in the said environment;
- (4) retrieving microorganisms from the said environment or a selected part thereof and isolating the nucleic acid from the retrieved microorganisms;
- (5) comparing any marker sequences in the nucleic acid isolated in step (4) to the unique marker sequence of each individual mutant stored as in step (2); and
- (6) selecting an individual mutant which does not contain any of the marker sequences as isolated in step (4).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2000:7170 USPATFULL

TITLE:

Identification of genes

INVENTOR(S):

Holden, David William, London, United Kingdom

PATENT ASSIGNEE(S):

Imperial College Innovations Limited, London, United

Kingdom (non-U.S. corporation)

NUMBER KIND DATE -----US 6015669 20000118 US 1997-871355 19970609 (8)

PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.:

Continuation of Ser. No. WO 1995-GB2875, filed on 11

Dec 1995

NUMBER DATE ______ GB 1994-24921 19941209 GB 1995-1881 19950131 GB 1995-9239 19950505 PRIORITY INFORMATION: WO 1995-GB2875 19951211 DOCUMENT TYPE: Utility

FILE SEGMENT:

Granted

PRIMARY EXAMINER: PRIMARY EXAMINER: Marschel, Ardin H. ASSISTANT EXAMINER: Whisenant, Ethan

LEGAL REPRESENTATIVE: Arnall Golden & Gregory, LLP

NUMBER OF CLAIMS: 26 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

116 Drawing Figure(s); 112 Drawing Page(s)

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L10 ANSWER 4 OF 12 USPATFULL on STN
- ΤI Identification of genes
- AB A method for identifying a microorganism having a reduced adaptation to a particular environment comprising the steps of:
 - (1) providing a plurality of microorganisms each of which is independently mutated by the insertional inactivation of a gene with a nucleic acid comprising a unique marker sequence so that each mutant contains a different marker sequence, or clones of the said microorganism;
 - (2) providing individually a stored sample of each mutant produced by step (1) and providing individually stored nucleic acid comprising the unique marker sequence from each individual mutant;
 - (3) introducing a plurality of mutants produced by step (1) into the said particular environment and allowing those microorganisms which are able to do so to grow in the said environment;

- (4) retrieving microorganisms from the said environment or a selected part thereof and isolating the nucleic acid from the retrieved microorganisms;
- (5) comparing any marker sequences in the nucleic acid isolated in step (4) to the unique marker sequence of each individual mutant stored as in step (2); and
- (6) selecting an individual mutant which does not contain any of the marker sequences as isolated in step (4).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

1999:27395 USPATFULL

TITLE:

Identification of genes

INVENTOR(S):

Holden, David William, London, United Kingdom

PATENT ASSIGNEE(S):

RPMS Technology Limited, London, United Kingdom

(non-U.S. corporation)

	NUMBER	KIND DATE	
PATENT INFORMATION:	US 5876931 WO 9617951	19990302 19760613	
APPLICATION INFO.:	US 1997-637759 WO 1995-GB2875	19970719 19951211	(8)
			PCT 371 date PCT 102(e) date

NUMBER									DATE													
	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	

PRIORITY INFORMATION:

GB 1994-24921 19941209 GB 1995-1881 19950131 GB 1995-9239 19950505

DOCUMENT TYPE:

Utility Granted

FILE SEGMENT: PRIMARY EXAMINER:

ASSISTANT EXAMINER:

Degen, Nancy Schwartzman, Robert

LEGAL REPRESENTATIVE: Arnall Golden & Gregory LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

31

NUMBER OF DRAWINGS:

119 Drawing Figure(s); 112 Drawing Page(s)

LINE COUNT:

6165

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L10 ANSWER 5 OF 12 USPATFULL on STN

Artificial chromosome vector TΤ

AB The present invention relates to a recombinant DNA molecule which contains the telomere and, optionally, the centromere of a higher eukaryote, particularly a plant, the telomere itself, the centromere itself, a method of producing a polypeptide in a recipient cell which utilizes said recombinant DNA molecule, host cells transformed with said recombinant molecule, and uses for said recombinant molecule.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

93:104847 USPATFULL

TITLE:

Artificial chromosome vector

INVENTOR(S):

Richards, Eric J., Lloyd Harbor, NY, United States Ausubel, Frederick M., Newton, MA, United States

PATENT ASSIGNEE(S):

The General Hospital Corporation, Boston, MA, United

States (U.S. corporation)

NUMBER	KIND	DATE
US 5270201		19931214

PATENT INFORMATION:

APPLICATION INFO.: US 1992-860585 19920330 (7)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1991-742554, filed on 9 Aug

1991, now abandoned which is a continuation of Ser. No.

US 1989-404525, filed on 8 Sep 1989, now abandoned which is a continuation-in-part of Ser. No. US 1988-172467, filed on 24 Mar 1988, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Schwartz, Richard A. ASSISTANT EXAMINER: Carter, Philip W.

LEGAL REPRESENTATIVE: Sterne, Kessler, Goldstein & Fox

NUMBER OF CLAIMS: 25 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 23 Drawing Figure(s); 19 Drawing Page(s)

LINE COUNT: 1901

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L10 ANSWER 6 OF 12 DGENE COPYRIGHT 2003 THOMSON DERWENT on STN

TI Yeast which ferments xylose to methanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated

at each of its multiple reiterated ribosomal

DNA sites

AN AAV12824 DNA DGENE

This sequence represents an amplification primer for the yeast 5S rDNA sequence. The amplified sequence can be used in the yeast of the invention, which ferments xylose to ethanol. The yeast comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple

reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol; the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at least 20 generations. The yeast is produced by integrating multiple copies of exogenous DNA into reiterated chromosomal DNA of cells. The yeast produced by the integration method, even upon culture in non-selective medium for multiple generations (e.g.

up to 20), retain their full capability to ferment xylose to ethanol.

ACCESSION NUMBER: AAV12824 DNA DGENE

TITLE: Yeast which ferments xylose to methanol - comprising xylitol

reductase, xylitol dehydrogenase and xylulokinase

66p

genes integrated at each of its multiple

reiterated ribosomal DNA sites

INVENTOR: Chen Z; Ho N W Y

PATENT ASSIGNEE: (PURD) PURDUE RES FOUND.
PATENT INFO: WO 9742307 A1 19971113

APPLICATION INFO: WO 1997-US7663 19970506 PRIORITY INFO: US 1996-16865 19960506

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: 1997-558974 [51]

DESCRIPTION: Primer for yeast 5S rDNA sequence.

L10 ANSWER 7 OF 12 DGENE COPYRIGHT 2003 THOMSON DERWENT on STN

Yeast which ferments xylose to methanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated at each of its multiple reiterated ribosomal

DNA sites

AN AAV12829 DNA DGENE

AB This sequence is an amplification primer for the yeast Tn903 kanamycin resistance gene. The amplified sequence can be used in the yeast of the

invention, which ferments xylose to ethanol. The yeast comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol; the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at least 20 generations. The yeast is produced by integrating multiple copies of exogenous DNA into reiterated chromosomal

even upon culture in non-selective medium for multiple generations (e.g. up to 20), retain their full capability to ferment xylose to ethanol.

ACCESSION NUMBER: AAV12829 DNA DGENE

TITLE: Yeast which ferments xylose to methanol - comprising xylitol

reductase, xylitol dehydrogenase and xylulokinase

genes integrated at each of its multiple

DNA of cells. The yeast produced by the integration method,

reiterated ribosomal DNA sites

INVENTOR: Chen Z; Ho N W Y

PATENT ASSIGNEE: (PURD) PURDUE RES FOUND.
PATENT INFO: WO 9742307 A1 19971113

APPLICATION INFO: WO 1997-US7663 19970506 PRIORITY INFO: US 1996-16865 19960506

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: 1997-558974 [51]

DESCRIPTION: Primer for yeast Tn 903 kanamycin resistance gene.

L10 ANSWER 8 OF 12 DGENE COPYRIGHT 2003 THOMSON DERWENT on STN

TI Yeast which ferments xylose to methanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated at each of its multiple reiterated ribosomal

DNA sites

AN AAV12828 DNA DGENE

This sequence is an amplification primer for the yeast Tn903 kanamycin resistance gene. The amplified sequence can be used in the yeast of the invention, which ferments xylose to ethanol. The yeast comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple

reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its

chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol; the yeasts of (b) and (c) retain their capacity for fermenting

xylose to ethanol when cultured under non-selective conditions for at least 20 generations. The yeast is produced by integrating multiple copies of exogenous DNA into reiterated chromosomal DNA of cells. The yeast produced by the integration method,

even upon culture in non-selective medium for multiple generations (e.g. up to 20), retain their full capability to ferment xylose to ethanol.

ACCESSION NUMBER: AAV12828 DNA DGENE

TITLE: Yeast which ferments xylose to methanol - comprising xylitol

reductase, xylitol dehydrogenase and xylulokinase

genes integrated at each of its multiple

reiterated ribosomal DNA sites

INVENTOR: Chen Z; Ho N W Y

PATENT ASSIGNEE: (PURD) PURDUE RES FOUND.
PATENT INFO: WO 9742307 A1 19971113
APPLICATION INFO: WO 1997-US7663 19970506

66p

66p

PRIORITY INFO: US 1996-16865 19960506

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: 1997-558974 [51]

DESCRIPTION: Primer for yeast Tn 903 kanamycin resistance gene.

L10 ANSWER 9 OF 12 DGENE COPYRIGHT 2003 THOMSON DERWENT on STN

Yeast which ferments xylose to methanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated

at each of its multiple reiterated ribosomal

DNA sites

AN AAV12827 DNA DGENE

This sequence is an amplification primer for the yeast Tn903 kanamycin resistance gene. The amplified sequence can be used in the yeast of the invention, which ferments xylose to ethanol. The yeast comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol; the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at least 20 generations. The yeast is produced by integrating

least 20 generations. The yeast is produced by integrating multiple copies of exogenous DNA into reiterated chromosomal DNA of cells. The yeast produced by the integration method, even upon culture in non-selective medium for multiple generations (e.g.

up to 20), retain their full capability to ferment xylose to ethanol.

ACCESSION NUMBER: AAV12827 DNA DGENE

TITLE: Yeast which ferments xylose to methanol - comprising xylitol

reductase, xylitol dehydrogenase and xylulokinase

66p

genes integrated at each of its multiple

reiterated ribosomal DNA sites

INVENTOR: Chen Z; Ho N W Y

PATENT ASSIGNEE: (PURD) PURDUE RES FOUND.
PATENT INFO: WO 9742307 A1 19971113

APPLICATION INFO: WO 1997-US7663 19970506 PRIORITY INFO: US 1996-16865 19960506

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: 1997-558974 [51]

DESCRIPTION: Primer for yeast Tn 903 kanamycin resistance gene.

L10 ANSWER 10 OF 12 DGENE COPYRIGHT 2003 THOMSON DERWENT on STN

Yeast which ferments xylose to methanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated at each of its multiple reiterated ribosomal DNA sites

AN AAV12826 DNA DGENE

This sequence is an amplification primer for the yeast Tn903 kanamycin resistance gene. The amplified sequence can be used in the yeast of the invention, which ferments xylose to ethanol. The yeast comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol; the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at

least 20 generations. The yeast is produced by integrating multiple copies of exogenous DNA into reiterated chromosomal DNA of cells. The yeast produced by the integration method,

even upon culture in non-selective medium for multiple generations (e.g. up to 20), retain their full capability to ferment xylose to ethanol.

ACCESSION NUMBER: AAV12826 DNA DGENE

TITLE: Yeast which ferments xylose to methanol - comprising xylitol

reductase, xylitol dehydrogenase and xylulokinase

66p

66p

genes integrated at each of its multiple

reiterated ribosomal DNA sites

INVENTOR: Chen Z; Ho N W Y

PATENT ASSIGNEE: (PURD) PURDUE RES FOUND.
PATENT INFO: WO 9742307 A1 19971113
APPLICATION INFO: WO 1997-US7663 19970506

APPLICATION INFO: WO 1997-US7663 19970506 PRIORITY INFO: US 1996-16865 19960506

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: 1997-558974 [51]

DESCRIPTION: Primer for yeast Tn 903 kanamycin resistance gene.

L10 ANSWER 11 OF 12 DGENE COPYRIGHT 2003 THOMSON DERWENT on STN

Yeast which ferments xylose to methanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated at each of its multiple reiterated ribosomal

DNA sites

AN AAV12825 DNA DGENE

AB This sequence represents an amplification primer for the yeast 5S rDNA sequence. The amplified sequence can be used in the yeast of the invention, which ferments xylose to ethanol. The yeast comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple reiterated ribosomal DNA sites; (b) multiple copies of exogenous DNA, including XR, XD, and XK genes, fused to non-qlucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol; the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at least 20 generations. The yeast is produced by integrating multiple copies of exogenous DNA into reiterated chromosomal DNA of cells. The yeast produced by the integration method, even upon culture in non-selective medium for multiple generations (e.g.

up to 20), retain their full capability to ferment xylose to ethanol.

ACCESSION NUMBER: AAV12825 DNA DGENE

TITLE: Yeast which ferments xylose to methanol - comprising xylitol

reductase, xylitol dehydrogenase and xylulokinase

genes integrated at each of its multiple

reiterated ribosomal DNA sites

INVENTOR: Chen Z; Ho N W Y

PATENT ASSIGNEE: (PURD) PURDUE RES FOUND.

PATENT INFO: WO 9742307 A1 19971113

APPLICATION INFO: WO 1997 US7663 19970506

APPLICATION INFO: WO 1997-US7663 19970506 PRIORITY INFO: US 1996-16865 19960506

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: 1997-558974 [51]

DESCRIPTION: Primer for yeast 5S rDNA sequence.

L10 ANSWER 12 OF 12 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

Yeast which ferments xylose to ethanol - comprising xylitol reductase, xylitol dehydrogenase and xylulokinase genes integrated at each of its multiple reiterated ribosomal DNA sites.

1997-558974 [51] WPIDS AN

least 20 generations.

AB 9742307 A UPAB: 19991020

Novel yeast which ferments xylose to ethanol, comprises: (a) xylose reductase (XR), xylitol dehydrogenase (XD) and xylulokinase (XK) genes integrated at each of its multiple

reiterated ribosomal DNA sites; (b) multiple

copies of exogenous DNA, including XR, XD, and XK genes, fused to non-glucose inhibited promoters integrated into its chromosomal DNA, where the yeast simultaneously ferments glucose and xylose to ethanol; or (c) multiple copies of an introduced DNA containing XR, XD and XK genes, where the yeast ferments xylose to ethanol, where the yeasts of (b) and (c) retain their capacity for fermenting xylose to ethanol when cultured under non-selective conditions for at

USE - The methods can produce yeast, which even upon culture in non-selective medium for multiple generations, e.g. up to 20, retain their full capability to ferment xylose to ethanol.

Dwq.0/12

ACCESSION NUMBER:

1997-558974 [51] WPIDS

DOC. NO. CPI:

C1997-178545

TITLE:

Yeast which ferments xylose to ethanol - comprising

xylitol reductase, xylitol dehydrogenase and xylulokinase

genes integrated at each of its multiple reiterated ribosomal

DNA sites.

DERWENT CLASS: INVENTOR(S):

D16 D17 E17 H06 CHEN, Z; HO, N W Y

PATENT ASSIGNEE(S):

(PURD) PURDUE RES FOUND

COUNTRY COUNT:

76

PATENT INFORMATION:

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A1 19971113 (199751) * EN WO 9742307 66

RW: AT BE CH DE DK EA ES FI FR GB GH GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE HU IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG US UZ VN YU

AU 9728301 A 19971126 (199813)

EP 898616 A1 19990303 (199913) EN

R: AT BE DE DK ES FI FR GB GR IE IT NL PT SE

CN 1225125 A 19990804 (199949)

JP 2000509988 W 20000808 (200043) 50

MX 9809223 A1 19990701 (200061)

AU 731102 B 20010322 (200122)

BR 9710963 A 20010731 (200146)

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PATENT NO K	IND	APPLICATION	DATE
WO 9742307	A1	WO 1997-US7663	19970506
AU 9728301 EP 898616	A Al	AU 1997-28301 EP 1997-922698	19970506 19970506
CN 1225125	A	WO 1997-US7663 CN 1997-196195	19970506 19970506
JP 2000509988	W	JP 1997-540153 WO 1997-US7663	19970506 19970506
MX 9809223	A1	MX 1998-9223	19981105
AU 731102	В	AU 1997-28301	19970506
BR 9710963	A	BR 1997-10963	19970506
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